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Control of sex-specific apoptosis in *C. elegans* by the BarH homeodomain protein CEH-30 and the transcriptional repressor UNC-37/Groucho

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Apoptosis is essential for proper development and tissue homeostasis in metazoans. It plays a critical role in generating sexual dimorphism by eliminating structures that are not needed in a specific sex. The molecular mechanisms that regulate sexually dimorphic apoptosis are poorly understood. Here we report the identification of the *ceh-30* gene as a key regulator of sex-specific apoptosis in *Caenorhabditis elegans*. Loss-of-function mutations in *ceh-30* cause the ectopic death of male-specific CEM neurons. *ceh-30* encodes a BarH homeodomain protein that acts downstream from the terminal sex determination gene *tra-1*, but upstream of, or in parallel to, the cell-death-initiating gene *egl-1* to protect CEM neurons from undergoing apoptosis in males. The second intron of the *ceh-30* gene contains two adjacent *cis*-elements that are binding sites for TRA-1A and a POU-type homeodomain protein UNC-86 and acts as a sensor to regulate proper specification of the CEM cell fate. Surprisingly, the N terminus of CEH-30 but not its homeodomain is critical for CEH-30's cell death inhibitory activity in CEMs and contains a conserved eh1/FIL domain that is important for the recruitment of the general transcriptional repressor UNC-37/Groucho. Our study suggests that *ceh-30* defines a critical checkpoint that integrates the sex determination signal TRA-1 and the cell fate determination and survival signal UNC-86 to control the sex-specific activation of the cell death program in CEMs through the general transcription repressor UNC-37.

[Keywords: *C. elegans*; apoptosis; sex-specific; *unc-37*/Groucho; *ceh-30*; transcriptional regulation]

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The development of sexually dimorphic traits, those systematic differences between two sexes of an organism, is an important process in metazoan development. In many cases, sex-specific apoptosis plays an essential role in shaping the sexual dimorphism of animals (Roberts et al. 1999). For example, during male development in mammals, a Mullerian-inhibiting substance triggers apoptosis and regression of the Mullerian ducts that would develop into the female reproductive system (Price et al. 1977). Alternately, the Wolffian ducts, which would mature into the male-specific reproductive structures, undergo apoptosis during the development of a female (Yin et al. 2006). Additional instances in which apoptosis regulates sexually dimorphic development include Bax-dependant sexual differentiation of cell numbers in two regions of

the mouse forebrain, loss of lactotrophs in male rats, and mammary epithelial cell destruction in male mice (Dunbar et al. 1999; Aoki et al. 2001; Forger et al. 2004). Apoptosis also plays important roles in some diseases with sexually dimorphic outcomes or treatment efficacies, such as susceptibility to diabetes and renal injury (Casteels et al. 1998; Blush et al. 2004). Studying the regulation of apoptosis leading to these sex-specific differences is important for our understanding of sexual dimorphism and its implications on the development and treatment of related human diseases.

Caenorhabditis elegans has two sexes, males and self-reproducing hermaphrodites, which also exhibit sexually dimorphic apoptosis. A pair of bilaterally symmetric motor neurons named HSNs (hermaphrodite-specific neurons) control egg-laying in hermaphrodites and undergo apoptosis in males, where they are not needed (Sulston and Horvitz 1977; Sulston et al. 1983). In contrast, four male-specific chemosensory neurons (CEMs; cephalic companion neurons), which are thought to mediate che-

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motaxis of the male toward the hermaphrodite during courtship behavior, undergo apoptosis in hermaphrodites (Sulston and Horvitz 1977; Sulston et al. 1983). The sex-specific deaths of HSNs and CEMs provide a simple model for studying the molecular mechanisms that control sexually dimorphic apoptosis.

During the development of *C. elegans* hermaphrodites, 131 cells die in an invariant pattern (Sulston and Horvitz 1977; Kimble and Hirsh 1979; Sulston et al. 1983). Genetic analysis in *C. elegans* has led to the identification of four genes (*egl-1*, *ced-9*, *ced-4*, and *ced-3*) that act in a sequential manner to induce apoptosis (Ellis and Horvitz 1986; Hengartner et al. 1992; Conradt and Horvitz 1998). Biochemical studies indicate that EGL-1, a BH3-only protein, induces cell death by binding to CED-9, a cell death inhibitor and human Bcl-2 homolog, causing the release of CED-4 from the CED-4/CED-9 complex tethered to the outer mitochondrial membrane and the subsequent CED-4-mediated activation of the cysteine protease CED-3 (the cell death executioner) (Hengartner et al. 1992; Conradt and Horvitz 1998; Parrish et al. 2000; Yan et al. 2005). How apoptosis is activated in specific cells in response to death signals is poorly understood.

The sexual development of *C. elegans* soma is controlled by a negative regulatory cascade, initiated by HER-1, a secreted protein, which promotes male development by inhibiting the activity of TRA-2, a transmembrane receptor (Kuwabara et al. 1992; Perry et al. 1993). TRA-2 acts by inhibiting the activities of three interacting cytoplasmic proteins—FEM-1, FEM-2, and FEM-3—which promote male development by inhibiting the activity of TRA-1A, the terminal global regulator of somatic sex determination, through an unknown mechanism (Doniach and Hodgkin 1984; Kimble et al. 1984; Hodgkin 1988; Chin-Sang and Spence 1996; Meyer 1997; Mehra et al. 1999). TRA-1A, a zinc-finger protein, promotes female development by transcriptionally activating female-specific genes and/or by repressing male-specific genes (Zarkower and Hodgkin 1992, 1993). One TRA-1A target is the death-activating gene *egl-1*, which contains a TRA-1A-binding site 5.6 kb downstream from its ORF (Conradt and Horvitz 1999). Several gain-of-function (*gf*) mutations in *egl-1* disrupt this TRA-1-binding site and cause ectopic expression of *egl-1* in hermaphrodite HSNs and inappropriate HSN death, suggesting that TRA-1A represses the expression of *egl-1* in hermaphrodite HSNs (Conradt and Horvitz 1999). However, as a general sex determination factor that affects many somatic cells in *C. elegans* (Zarkower and Hodgkin 1992), TRA-1A does not affect most of the somatic cell deaths and needs to act with HSN-specific factors to control the sexually dimorphic death of HSNs.

Far less is known about what controls the life versus death decision of the male-specific CEM neurons. We undertook a genetic screen to search for mutations affecting CEM cell fates and sex-specific apoptosis. Here, we report the characterization of *ceh-30*, which is required specifically for the survival of CEMs in males and encodes a BarH homeodomain protein. We show that

TRA-1A and a POU-type homeodomain protein, UNC-86, act concertedly to regulate sex-specific expression of *ceh-30* in CEMs, leading to proper CEM cell death specification. CEH-30 then acts through a conserved eh1/FIL domain at its N terminus to recruit the general transcriptional repressor Groucho/UNC-37, resulting in the formation of a transcriptional repressosome that suppresses programmed cell death. Thus *ceh-30* represents a critical link between the sex determination pathway and the programmed cell death pathway that controls sex-specific apoptosis in *C. elegans*.

Results

A genetic screen to identify genes that regulate CEM cell death specification

To identify factors that control sex-specific CEM death, we carried out a green fluorescent protein (GFP)-based screen using a CEM-specific reporter construct ($P_{pkd-2}gfp$) to isolate mutations that alter the sexually dimorphic cell fates of CEMs (Barr and Sternberg 1999). The *pkd-2* promoter drives GFP expression in four CEM neurons and some ray neurons in the tail of male animals; no GFP expression is observed in hermaphrodites (data not shown). We generated an integrated transgenic array containing $P_{pkd-2}gfp$ (*smIs23*) in *him-5(e1490)* animals, which produce a high frequency of male progeny as a result of increased meiotic nondisjunction (Goldstein 1986). We then performed ethyl methane sulfonate (EMS) mutagenesis on the resulting strain, *smIs23; him-5(e1490)*, and screened for mutations that cause ectopic CEM loss in males or improper CEM survival in hermaphrodites (see Materials and Methods).

We screened 8500 haploid genomes and isolated several mutations that affect the cell fate specification of CEMs and/or HSNs (E. Peden, E. Kimberly, and D. Xue, unpubl.), including the *sm130* mutation and a mutation (*sm117*) that affects a previously identified gene, *unc-86*.

sm130 is a X-linked mutation that causes ectopic CEM death in males

In *him-5(e1490)* animals carrying *smIs26*, another $P_{pkd-2}gfp$ -containing integrated array that also harbors $P_{tph-1}gfp$ (a HSN reporter) (Sze et al. 2000), 100% of CEMs survive in males (Table 1). In contrast, only 30% of CEMs are present in *smIs26; him-5(e1490); ceh-30(sm130)* males. The loss of CEMs in *sm130* males can be suppressed by a strong loss-of-function mutation in either the *egl-1* gene (*n3082*) or the *ced-3* gene (*n717*), suggesting that these CEMs die by programmed cell death rather than adopting a non-CEM cell fate (Table 2). *sm130* does not affect the sex-specific death of HSNs (Table 1) or the deaths and sexual differentiation of other somatic cells (data not shown). Therefore, *sm130* appears to affect a gene specifically regulating CEM cell death.

Using the phenotype of ectopic male CEM death, *sm130* was mapped to Linkage Group X (LGX) between

Table 1. *ceh-30* alleles cause specific loss of CEMs in males

| Genotype | % CEMs surviving (<i>n</i>) | | % HSNs surviving (<i>n</i>) ^a | |
|-----------------------|-------------------------------|-----------|--|--------|
| | Hermaphrodite | Male | Hermaphrodite | Male |
| Wild type | 0 (104) | 100 (100) | 100 (40) | 0 (16) |
| <i>ceh-30(sm130)</i> | 0 (80) | 30 (116) | 98 (62) | 0 (50) |
| <i>ceh-30(tm272)</i> | 0 (100) | 0 (100) | 100 (50) | 0 (50) |
| <i>ceh-30(tm2157)</i> | 0 (100) | 71 (100) | 100 (50) | 0 (50) |

^aThe presence of CEMs and HSNs was scored as described in Materials and Methods.

The complete genotype of the animals scored was, from top to bottom, as follows: *smIs26; him-5(e1490)*, *smIs26; him-5(e1490); ceh-30(sm130)*, *smIs26; him-5(e1490); ceh-30(tm272)*, and *smIs26; him-5(e1490); ceh-30(tm2157)*.

n indicates the number of presumptive CEMs or HSNs scored.

two genetic markers, *lon-2* and *unc-2*, at ~3.03 Mb or ~12.7 cM, which was covered by three overlapping Cosmids, M02F4, C33D12, and F52E4 (see Fig. 1A; Materials and Methods).

ceh-30 encodes a BarH homeodomain protein that is essential for CEM survival in males

To clone the gene affected by *sm130*, germline transformations of *smIs23; him-5(e1490); sm130* animals were carried out by injecting three individual Cosmids (M02F4, C33D12, and F52E4) spanning the locus defined by the *sm130* mutation. Only Cosmid C33D12 showed rescue of the ectopic male CEM death phenotype. Further analysis of C33D12 subclones indicates that a single ORF, *ceh-30*, which encodes a BarH homeodomain-containing protein, is responsible for the rescuing activity (Fig. 1A).

Sequencing of genomic DNA from the *sm130* mutant revealed a single nucleotide substitution (a C-to-T transition) 336 base pairs (bp) into the second intron of *ceh-30* (Fig. 1A). We also obtained two deletion alleles of *ceh-30*, *tm272*, and *tm2157*, which remove 425 bp and 718 bp from the *ceh-30* coding region, respectively (Fig. 1A). *ceh-30(tm272)* deletes part of the second intron of *ceh-30*, including the nucleotide mutated by *sm130*, and causes 100% CEM death in mutant males (Table 1). *ceh-30(tm2157)* removes part of exon 1 and all of intron 1 and exon 2 but leaves most of intron 2 intact, including the region affected by *sm130* (Fig. 1A). Interestingly, *tm2157* only causes a weak reduction of male CEM survival (29%), albeit removing most of the CEH-30 protein including its entire homeodomain (see below). These results underscore the critical role of the *ceh-30* second intron in protecting CEMs from undergoing apoptosis in males. Like *sm130*, *ceh-30(tm272)* and *ceh-30(tm2157)* do not affect the death of HSNs or the death and sex differentiation of other somatic cells (Table 1; data not shown).

ceh-30 acts downstream from *tra-1* and upstream of, or in parallel to, *egl-1* to regulate sex-specific CEM death

To understand how *ceh-30* acts to control sexually dimorphic apoptosis of CEMs, we carried out genetic epistasis analysis to determine the relationships between

ceh-30 and key regulators of sex determination and apoptosis. In particular, we focused on the terminal sex determination gene *tra-1*, the cell death initiator *egl-1*, and the cell death executor *ced-3* (Ellis and Horvitz 1986; Zarkower and Hodgkin 1992; Conradt and Horvitz 1999).

tra-1 acts at the terminal step of the *C. elegans* sex determination pathway to control the sexual fates of somatic cells, including the fates of HSN and CEM neurons. Hermaphrodite XX animals homozygous for the null *tra-1(e1099)* mutation develop as low-fertility males with 100% inappropriately surviving CEMs (Table 2; Hodgkin and Brenner 1977). However, in *smIs23; tra-1(e1099); ceh-30(tm272)* XX animals, CEMs are never present. Given that *tra-1* is the terminal sex determination gene, this result suggests that *ceh-30* acts downstream from, or in parallel to, *tra-1* as well as the entire sex determination pathway to control the sex-specific death of CEMs.

Table 2. *ceh-30* acts downstream from *tra-1* and upstream of, or in parallel to, *egl-1* to regulate sex-specific CEM death

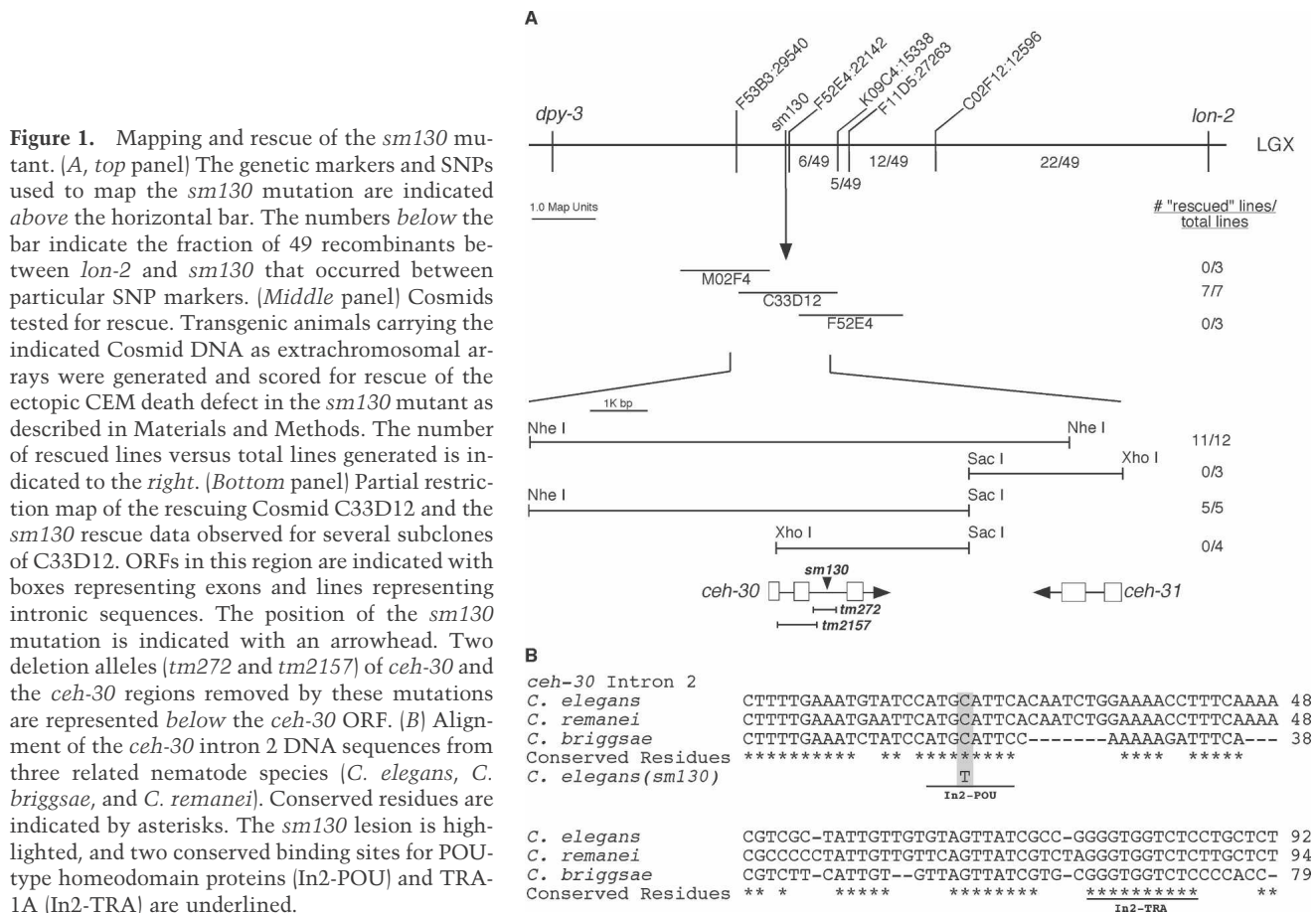
| Genotype | % CEMs surviving (<i>n</i>) ^a | |
|------------------------------------|--|-----------|
| | Hermaphrodite | Male |
| Wild type | 0 (100) | 100 (104) |
| <i>ceh-30(tm272)</i> | 0 (100) | 0 (100) |
| <i>tra-1(e1099)</i> | 100 (100) | ND |
| <i>tra-1(e1099); ceh-30(tm272)</i> | 0 (100) | ND |
| <i>egl-1(n3082)</i> | 53 (80) | 100 (80) |
| <i>egl-1(n3082)/itDf2</i> | 90 (98) | ND |
| <i>egl-1(n3082); ceh-30(tm272)</i> | 54 (80) | 94 (80) |
| <i>ced-3(n717)</i> | 85 (80) | 98 (80) |
| <i>ced-3(n717); ceh-30(tm272)</i> | 79 (80) | 98 (80) |

^aThe presence of CEMs was scored as described in Materials and Methods.

The complete genotype of the animals scored was, from top to bottom, as follows: *smIs23; him-5(e1490)*, *smIs23; him-5(e1490); ceh-30(tm272)*, *smIs23; tra-1(e1099) dpy-18(e364)*, *smIs23; tra-1(e1099) dpy-18(e364); ceh-30(tm272)*, *smIs23; him-8(e1489); egl-1(n3082)*, *smIs23/+; him-8/+; egl-1(n3082)/itDf2*, *smIs23; him-8(e1489); egl-1(n3082); ceh-30(tm272)*, *smIs23; ced-3(n717); him-5(e1490)*, and *smIs23; ced-3(n717); him-5(e1490); ceh-30(tm272)*.

(ND) Not determined.

n indicates the number of presumptive CEMs scored. *itDf2* is a deficiency that deletes the *egl-1* locus.



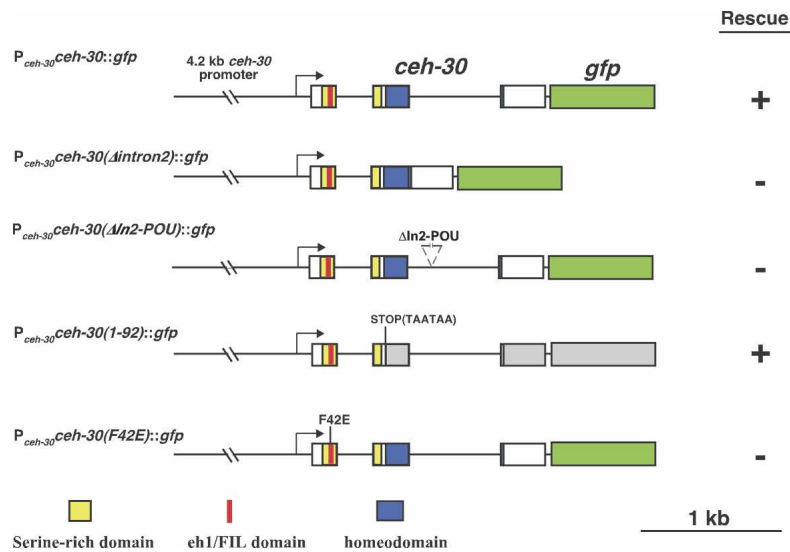
egl-1 is proposed to initiate the activation of all somatic cell deaths in *C. elegans* (Conradt and Horvitz 1998). In *sm130*; *egl-1*(n3082); *ceh-30*(*tm272*) males, the ectopic CEM death phenotype caused by *ceh-30*(*tm272*) is almost completely suppressed by *egl-1*(n3082), suggesting that *ceh-30* acts upstream of, or in parallel to, *egl-1* to control sex-specific CEM deaths. Intriguingly, *ceh-30* can also regulate CEM death in the absence of either *egl-1* or *ced-9*, suggesting that *ceh-30* can act in parallel to *egl-1* and *ced-9* to control sex-specific CEM death (H.T. Schwartz and H.R. Horvitz, pers. comm.). Like *egl-1*(n3082), *ced-3*(n717) also potently suppresses ectopic CEM death in *ceh-30*(*tm272*) males (Table 2). Taken together, these results suggest that *ceh-30* acts downstream from the sex determination pathway but upstream of, or in parallel to, the *C. elegans* programmed cell death pathway to control sexually dimorphic deaths of CEM neurons and may serve as a critical mediator between these two globally acting pathways to regulate cell-type and sex-specific deaths.

ceh-30(*sm130*) and *ceh-30*(*tm272*) affect a conserved binding site for POU-type homeodomain proteins

Two *ceh-30* alleles, *sm130* and *tm272*, alter or remove sequences within intron 2 of the *ceh-30* gene. The C-to-T transition caused by *sm130* falls within a conserved se-

quence motif (CATGCATTC) in intron 2 of the *ceh-30* genes from three different nematode species (*Caenorhabditis briggsae*, *Caenorhabditis remanei*, and *Caenorhabditis elegans*) (Fig. 1B). By searching the Transcription Factor Database (TFD), we found that this sequence shares high sequence identity with the binding sites of the mammalian POU-type homeodomain proteins GHF/Pit1 (ATGCATTC) and Brn-2/N-Oct-3 [ATG(A/C)AT(A/T)] (Argenton et al. 1996; Rhee et al. 1998). In addition, the *C. elegans* POU-type homeodomain protein UNC-86 has been shown to recognize a similar sequence, ATG(A/C)AT (Finney et al. 1988; Xue et al. 1992). These observations suggest that *sm130* might affect the binding site of a POU-type homeodomain protein and that this site, which we call In2-POU, is important for *ceh-30*'s function in preventing CEM death in males.

There are several additional conserved sequence motifs in intron 2 downstream from In2-POU (Fig. 1B). One of them, located 55 bp downstream, resembles a conserved TRA-1A-binding site (Zarkower and Hodgkin 1993). This site, which we name In2-TRA, is altered in several *ceh-30* gain-of-function mutants where CEMs survive inappropriately in hermaphrodites (H.T. Schwartz and H.R. Horvitz, pers. comm.). These findings suggest that intron 2 of *ceh-30* is critical for the regulation of sex-specific CEM death. Indeed, the *tm272* dele-



Rescue

Figure 2. The N terminus of CEH-30 but not its homeodomain is required for its activity to protect male CEMs from cell death. Graphical presentations of five *cek-30* constructs are shown. Boxes indicate *cek-30* exons or the coding region for GFP (in green). The region encoding the homeodomain is highlighted with blue, the region encoding the SRD is highlighted with yellow, and the region encoding the eh1/FIL motif is highlighted with red. Transgenic *smIs26*; *him-5(e1490)*; *cek-30(tm272)* animals carrying the indicated DNA construct as extrachromosomal arrays were scored for rescue of the ectopic CEM death defect as described in Materials and Methods. For each construct, at least three independent transgenic lines were scored and found to have similar results. (+) Rescue; (–) failure to rescue.

tion that removes both In2-POU and In2-TRA motifs results in a complete absence of CEMs in males.

To verify the importance of *cek-30* intron 2 and In2-POU in regulating the sexually dimorphic cell fate of CEMs, we generated a *cek-30* translational GFP fusion (*P_{cek-30}cek-30::gfp*) that contains a 4.2-kb *cek-30* promoter as well as the entire *cek-30* ORF and found that this construct rescues the CEM defect of the *cek-30(tm272)* mutant (Fig. 2). In contrast, a *P_{cek-30}cek-30(Δintron2)::gfp* construct that lacks all of intron 2 or a *P_{cek-30}cek-30(ΔIn2-POU)::gfp* construct in which the 8-bp In2-POU site (ATGCATTC) is deleted failed to rescue the *cek-30(tm272)* mutant (Fig. 2), providing further evidence that intron 2 and the In2-POU site of *cek-30* are critical for its apoptosis-inhibitory activity in male CEMs.

unc-86 is required for CEM cell fate specification independent of *ced-3*

Our CEM cell fate mutant screen also identified an allele of *unc-86* (*sm117*) that results in almost complete absence of GFP expression at the CEM positions in *smIs23* males (Table 3). However, very few GFP-expressing cells are seen at the CEM positions in *smIs23*; *unc-86(sm117)*; *ced-3(n717)* males, indicating that *unc-86* is required for CEM cell fate specification at a step prior to the life versus death decision of CEMs. We also observed almost complete loss of GFP expression at the CEM positions in several other *unc-86* loss-of-function mutants (Table 3). UNC-86 is expressed in 47 neurons during *C. elegans* development, including male CEMs and hermaphrodite HSNs (Finney and Ruvkun 1990), and is critical for the proper differentiation of these UNC-86-expressing neurons, including the differentiation and survival of CEM neurons in males (Finney et al. 1988; Shaham and Bargmann 2002). Interestingly, the mouse ortholog of UNC-86, Brn3c, has been shown to act genetically upstream of the mouse BarH homeodomain protein Barhl1 to pro-

mote terminal differentiation and survival of mechanosensory hair cells (Li et al. 2002), suggesting a conserved regulatory relationship between these two types of homeodomain proteins. Therefore, *unc-86* is an excellent candidate for the transcriptional activator of *cek-30* through the In2-POU *cis*-element.

In addition to *unc-86*, there are only two other genes in the *C. elegans* genome that encode POU-type homeodomain proteins (*cek-18* and *cek-6*). The *cek-18*-null allele (*mg57*) does not cause any defect in CEM cell fate specification (Table 3; Greenstein et al. 1994). Animals homozygous for the only available *cek-6* allele (*mg60*) are lethal at the embryonic and L1 larval stages, making analysis of the CEM cell fate impossible. Analysis of CEH-6 expression using anti-CEH-6 antibody and reporter constructs does not identify CEH-6 expression in CEMs or HSNs (Burglin and Ruvkun 2001). These data

Table 3. *unc-86* affects CEM cell fate

| Genotype | % Male CEMs or <i>pkd-2</i> -positive cells present (n) ^a |
|-----------------------------------|--|
| Wild type | 100 (100) |
| <i>unc-86(sm117)</i> | 3 (80) |
| <i>unc-86(sm117); ced-3(n717)</i> | 1 (120) |
| <i>unc-86(e1416)</i> | 3 (80) |
| <i>unc-86(e1507)</i> | 4 (80) |
| <i>unc-86(n845)</i> | 4 (80) |
| Wild type | 100 (100) |
| <i>cek-18(mg57)</i> | 99 (332) |

^aThe presence of CEMs or *pkd-2*-positive cells was scored as described in Materials and Methods.

The complete genotype of the animals scored was, from top to bottom, as follows: *smIs23*; *him-5(e1490)*, *smIs23*; *unc-86(sm117)*; *him-5(e1490)*, *smIs23*; *unc-86(sm117); ced-3(n717)*; *him-5(e1490)*, *smIs23*; *unc-86(e1416)*; *him-5(e1490)*, *smIs23*; *unc-86(e1507)*; *him-5(e1490)*, *smIs23*; *unc-86(n845)*; *him-5(e1490)*, *smIs26*; *him-5(e1490)*, and *smIs26*; *him-5(e1490)*; *cek-18(mg57)*.

n indicates the number of CEMs or CEM-like cells scored.

suggest that *ceh-18* does not play a role in CEM cell fate specification, while a role for *ceh-6* in CEM fate determination is unlikely but cannot be ruled out.

UNC-86 and TRA-1A bind intron 2 of *ceh-30* in vitro

We next used the gel mobility shift assay to investigate the possibility that UNC-86 binds the In2-POU site. We found that full-length UNC-86 protein tagged with glutathione S-transferase (GST) at its N terminus (GST::UNC-86) can bind to a known UNC-86-binding site (CS2) from the promoter of the *mec-3* gene (Xue et al. 1993) as well as a 33-bp DNA fragment containing either the In2-POU site or the altered In2-POU site [In2-POU(m)] found in the *sm130* mutant (Fig. 3A). Quantification of the amount of DNA shifted due to GST::UNC-86 binding in four independent experiments reveals that binding to In2-POU(m) (12% DNA shifted) is significantly weaker than binding to In2-POU (28% DNA shifted; P -value < 0.0005) (Fig. 3B). This decrease in binding efficiency of GST::UNC-86 to the In2-POU(m) site is consistent with the reduced, but not abolished, *ceh-30* activity observed in the *ceh-30(sm130)* mutant (Table 1).

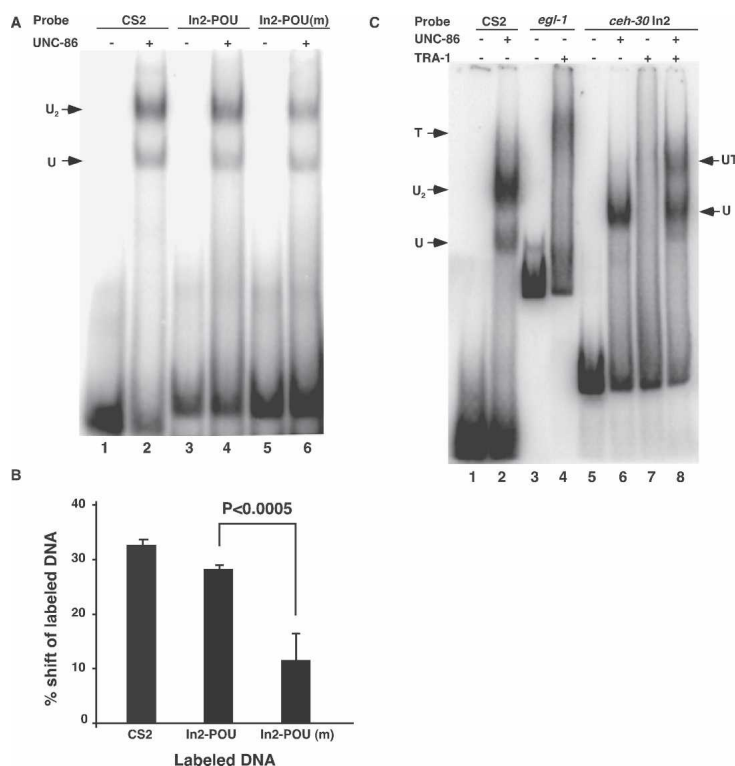
Given that *tra-1* acts upstream of *ceh-30* and that a potential TRA-1A-binding site (In2-TRA) is located 55 bp downstream from the In2-POU site, we investigated if TRA-1A binds to this site and if the binding of TRA-1A to this site affects the binding of UNC-86 to the neighboring In2-POU site or vice versa. As shown in Figure 3C, the full-length TRA-1A protein tagged with hexa-

histidine (TRA-1::His₆) can bind a 257-bp DNA fragment containing the critical TRA-1A-binding site from the *egl-1* gene (Fig. 3C, lanes 3,4; Conradt and Horvitz 1999). When we included a 91-bp DNA fragment carrying both the In2-POU site and the In2-TRA site in the binding reactions, we found that both GST::UNC-86 and TRA-1::His₆ were capable of binding to this DNA fragment, although the binding of TRA-1A to this DNA fragment was less stable, displaying a smear gel-shift pattern reflective of the formation of unstable DNA/protein complexes (Fig. 3C, lanes 5–7). Interestingly, addition of GST::UNC-86 to the TRA-1A-binding reaction stabilizes the binding of TRA-1::His₆ to this 91-bp *ceh-30* intronic sequence, resulting in the formation of a discrete TRA-1/UNC-86 supershifted band at the expense of the UNC-86/DNA complexes (Fig. 3C, lane 8). We confirmed the presence of both TRA-1 and UNC-86 in this DNA complex by an antibody supershift experiment (Supplementary Fig. S1). The observation that UNC-86 can stabilize the binding of TRA-1A to the intron 2 sequence suggests that these two proteins interact on this important DNA element to regulate the expression of *ceh-30* and thus the proper sex-specific death of CEM neurons.

CEH-30 is widely expressed during embryonic development including in male CEM neurons

We next determined the expression pattern of *ceh-30* during *C. elegans* development using an integrated transgene (*sm1s54*) carrying $P_{ceh-30}ceh-30::gfp$ that rescues

Figure 3. Both UNC-86 and TRA-1A bind to the intron 2 sequence of *ceh-30* in vitro. (A) Binding of GST::UNC-86 to a POU-type homeodomain-binding site (In2-POU) found in intron 2 of *ceh-30* and an altered site [In2-POU(m)] derived from the *sm130* mutant. Twenty-five nanograms of recombinant GST::UNC-86 (lanes 2,4,6) or binding buffer alone (lanes 1,3,5) were incubated with labeled oligonucleotides derived from a known UNC-86-binding site (CS2; lanes 1,2), In2-POU (lanes 3,4), or In2-POU(m) (lanes 5,6) and resolved on an 8% native polyacrylamide gel. (U) The gel shift species containing the UNC-86 monomer; (U₂) the gel shift species containing the UNC-86 homodimer (Xue et al. 1993). (B) The *sm130* lesion reduces the binding of GST::UNC-86 to In2-POU. The fraction of labeled probe shifted by GST::UNC-86 was quantified from four independent experiments as described in Materials and Methods. (C) UNC-86 stabilizes the binding of TRA-1A to the intron 2 sequence of *ceh-30*. Twenty-five nanograms of recombinant GST::UNC-86 and 5 ng of TRA-1::His₆ either alone or together were incubated with labeled CS2 (lanes 1,2), 257-bp DNA fragment containing the TRA-1A-binding site from the *egl-1* gene (*egl-1*, lanes 3,4), or 91-bp intron 2 DNA fragment from *ceh-30* containing both In2-POU and In2-TRA (*ceh-30* In2, lanes 5–8). The reactions were resolved on a 4% native polyacrylamide gel. (T) The gel shift species containing the TRA-1 protein; (UT) the gel shift species containing both UNC-86 and TRA-1A.



the CEM defect of the *ceh-30(tm272)* mutant (Fig. 2). The CEH-30::GFP fusion is expressed widely during embryonic development, and the expression is especially robust in the anterior half of the embryos (Fig. 4A). CEH-30::GFP expression continues into the larval and adult stages, albeit at greatly reduced levels.

We then examined whether CEH-30 is expressed in the CEM neurons by following the expression of CEH-30::GFP in ventral CEM neurons at ~400–430 min after first cleavage, which are slightly after their births at ~320 min (Sulston et al. 1983). In hermaphrodite *smIs54* embryos, CEH-30::GFP was never seen in ventral CEMs (0/15 embryos) (Fig. 4C); this result is consistent with the predicted absence of CEH-30 expression in hermaphrodite CEMs. In contrast, in *tra-1(n1099); smIs54* XX embryos, which are completely masculinized and virtually identical to male embryos, CEH-30::GFP was observed in all ventral CEMs (15/15 embryos) (Fig. 4B,C), suggesting that CEH-30 is expressed in developing male CEM neurons. Given that TRA-1A directly binds to intron 2 of *ceh-30* (Fig. 3C) and mutations in In2-TRA of *ceh-30* cause improper CEM survival in hermaphrodites (H.T. Schwartz and H.R. Horvitz, pers. comm.), this result suggests that TRA-1 directly suppresses CEH-30 expression in hermaphrodite CEMs. Interestingly, in *unc-86(e1416) tra-1(e1099); smIs54* XX masculinized embryos, CEH-

30::GFP was not seen in ventral CEM-like cells (0/15 embryos) (Fig. 4C), suggesting that UNC-86 is important for CEH-30 expression in CEMs. Taken together, these results are consistent with the model that UNC-86 activates and TRA-1 inhibits the expression of CEH-30 in CEMs.

The CEH-30 homeodomain is largely dispensable for CEM survival in males

Homeodomains mediate binding to DNA and are critical for the functions of homeodomain-containing proteins (Hayashi and Scott 1990). One *ceh-30* deletion allele, *tm2157*, removes the last 45 bp of exon 1, the entirety of intron 1 and exon 2, and the first 141 bp of intron 2, but leaves the In2-POU site and the In2-TRA site intact (Fig. 1A). This deletion results in the removal of the CEH-30 homeodomain, and presumably, the synthesis of a truncated CEH-30 protein containing only the first 46 amino acids. Despite the large truncation, *ceh-30(tm2157)* animals show only a weak reduction (29% reduction) in male CEM survival (Table 1), suggesting that the homeodomain of CEH-30 is largely dispensable for its activity in protecting against CEM cell death in males. To confirm this result, we generated a *ceh-30::gfp* fusion construct that carries tandem stop codons at the begin-

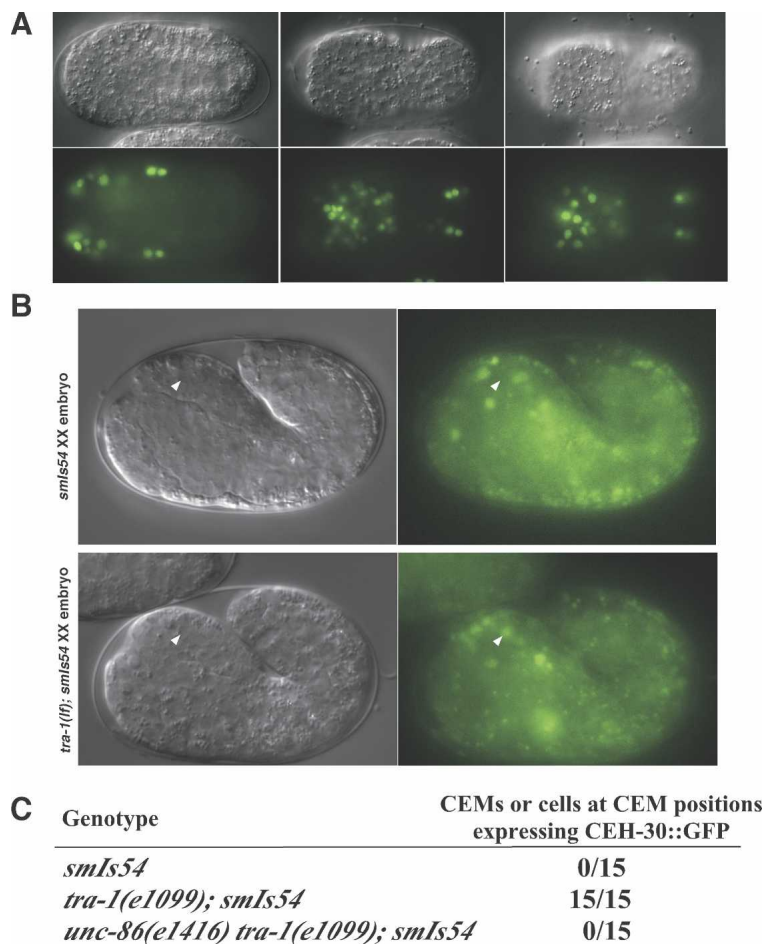


Figure 4. CEH-30 expression patterns in *C. elegans* embryos. (A) GFP expression directed from an integrated array (*smIs54*) containing $P_{ceh-30}ceh-30::gfp$ at ~360 min post-first cleavage. Three different focal planes of the same embryo are shown with the anterior toward the left and posterior toward the right. (B) Expression of CEH-30::GFP in a *smIs54* XX embryo (~420 min post-first cleavage, top panel) and a *tra-1(e1099); smIs54* XX embryo at the similar stage (bottom panel). The ventral CEM neuron is indicated by a white arrowhead. (C) Frequency of CEH-30::GFP expression in ventral CEMs of XX embryos at ~380–430 min after first cleavage. The genotypes of XX embryos are indicated.

ning of the *ceh-30* BarH homeodomain-coding region [*P_{ceh-30}ceh-30(1–92)::gfp*], resulting in the synthesis of a 92-amino-acid N-terminal fragment of CEH-30. This construct rescues the *ceh-30(tm272)* defect (Fig. 2), providing further support to the idea that the N terminus of CEH-30 is capable of specifying the CEM life versus death decision in the absence of its homeodomain.

The N-terminal eh1/FIL motif of CEH-30 and UNC-37/Groucho are required to protect against CEM cell death in males

Analysis of the CEH-30 protein sequence reveals only three conserved domain structures: the BarH homeodomain (amino acids 94–154), a serine-rich domain (SRD) (amino acids 29–80), and an engrailed homology domain or FIL (eh1/FIL) protein interaction motif (amino acids 42–49) embedded within the SRD (Fig. 5A). The putative *ceh-30* transcript made in the *ceh-30(tm2157)* mutant would encode a truncated CEH-30 protein with only the first 17 amino acids of the SRD domain (52 amino acids in length) and the first five amino acids of the eh1/FIL motif (FRISD from FRISDILE). The eh1/FIL domain was first identified as a motif required for the association of fly homeodomain proteins with the general transcriptional repressor *Groucho* (Smith and Jaynes 1996; Choi et al. 1999; Jimenez et al. 1999; Bae et al. 2003).

We thus tested whether this motif is important for protecting against CEM death in males by substituting Phe 42, a conserved and critical residue in the eh1/FIL motif, with glutamate (Smith and Jaynes 1996; Jennings et al. 2006). A *ceh-30::gfp* fusion construct harboring this mutation [*P_{ceh-30}ceh-30(F42E)::gfp*] failed to rescue the *ceh-30(tm272)* mutant, indicating that the eh1/FIL motif of CEH-30 is required to inhibit CEM cell death in males (Fig. 2).

We then examined animals defective in the *unc-37* gene, which encodes the *C. elegans* *Groucho* ortholog (Pflugrad et al. 1997; Winnier et al. 1999), to determine if *unc-37* is involved in CEM cell fate specification. Male animals carrying the only viable allele of *unc-37(e262)* showed decreased CEM survival (Table 4), whereas the *unc-37(e262); ced-3(n2433)* double mutant showed normal CEM survival in male animals. We also examined CEM survival in *unc-37(e262); ceh-30(lf)* double mutants and found that two partial loss-of-function *ceh-30* alleles (*sm130* and *tm2157*) either additively or synergistically reduced CEM survival with the *unc-37(e262)* mutation (Table 4). In particular, *unc-37(e262)* and *ceh-30(tm2157)* mutations, which individually only mildly reduce male CEM survival (17% and 29% male CEM death, respectively), can synergistically cause most CEMs to die in males (74% CEM death). These results indicate that *unc-37* is important for maintaining CEM survival in males and likely acts with CEH-30 to inhibit male CEM cell death.

Finally, we performed yeast two-hybrid experiments to examine if CEH-30 interacts with UNC-37 and if the eh1/FIL motif of CEH-30 mediates the interaction of CEH-30 with UNC-37. In these assays, we fused either CEH-30 or UNC-37 to the GAL4 activation domain (AD) or DNA-binding domain (DB) and tested if the fusion proteins interact in yeast to induce the synthesis of the HIS3 and URA3 enzymes as well as the synthesis of β -D-galactosidase (lacZ). We found that yeast strains expressing both GAL4-AD-CEH-30 and GAL4-DB-UNC-37 or expressing both GAL4-DB-CEH-30 and GAL4-AD-UNC-37 grew on selective plates lacking histidine and containing 10 mM 3-amino-1,2,4-triazole (3AT, an inhibitor of the HIS3 enzyme), failed to grow on selective plates with 0.2% 5-fluoroorotic acid (5FOA, which is converted to toxic 5-fluorouracil by URA3), and had significantly

Figure 5. CEH-30 and UNC-37 interact through the eh1/FIL motif. (A) Alignment of the eh1/FIL motifs in various proteins from *C. elegans*, *Drosophila* (*D. Mel*), mouse (*M. Mus*), and human (*H. sap*). Highly conserved amino acid residues are highlighted. (B) Yeast two-hybrid assays. (Panel 1) Yeast transformants expressing CEH-30 and UNC-37 GAL4 fusions as indicated were spotted on synthetic complete medium lacking tryptophan and leucine (SC-Leu-Trp). (Panel 2) Growth on the SC-Leu-Trp-His plate with 10 mM 3AT. Growth indicates an interaction between the two tested fusion proteins. (Panel 3) Growth on the SC-Leu-Trp-Ura plate containing 0.2% 5FOA. Poor or no growth on the SC-Leu-Trp-Ura plate with 0.2% 5FOA indicates interaction between two tested fusion proteins. (Panel 4) β -Galactosidase staining was performed on a nitrocellulose membrane as described in Materials and Methods. Dark staining indicates interaction between the two tested fusion proteins. β -Galactosidase activity was also measured by the Chlorophenol red- β -D-galactopyranoside (CPRG) cleavage assay and is indicated below the staining assay. The numbers shown are folds of increase in activity over the yeast strain that expressed GAL4-AD and GAL4-DB. Full-length CEH-30 (wild type or F42E mutant) and UNC-37 proteins were fused to either the GAL4-AD or the GAL4-DB. GAL4-DB/GAL4-AD and GAL4-DB::E2F1/GAL4-AD::RB were used as negative and positive controls for protein interaction, respectively.

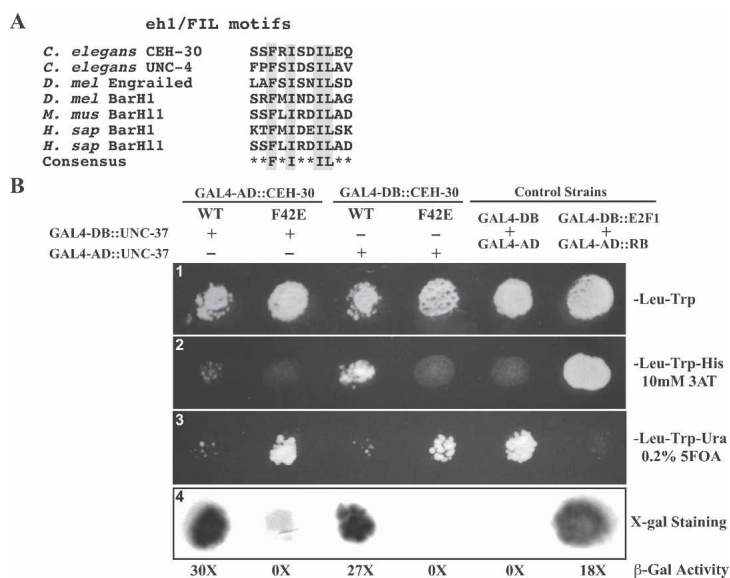


Table 4. *unc-37* affects CEM cell fate

| Genotype | % Male CEMs surviving (n) ^a |
|-------------------------------------|--|
| Wild type | 100 (100) |
| <i>unc-37(e262)</i> | 83 (496) |
| <i>unc-37(e262); ced-3(n2433)</i> | 99 (260) |
| <i>ceh-30(sm130)</i> | 30 (116) |
| <i>unc-37(e262); ceh-30(sm130)</i> | 14 (208) |
| <i>ceh-30(tm2157)</i> | 71 (100) |
| <i>unc-37(e262); ceh-30(tm2157)</i> | 26 (208) |
| <i>ceh-30(tm272)</i> | 0 (100) |
| <i>unc-37(e262); ceh-30(tm272)</i> | 0 (80) |

^aThe presence of CEMs was scored as described in Materials and Methods.

The complete genotype of the animals scored was, from top to bottom, as follows: *smIs26; him-5(e1490); bli-4(e937) unc-37(e262); smIs26; him-5(e1490); bli-4(e937) unc-37(e262); smIs26 ced-3(n2433); him-5(e1490); smIs26; him-5(e1490); ceh-30(sm130); bli-4(e937) unc-37(e262); smIs26; him-5(e1490); ceh-30(sm130); smIs26; him-5(e1490); ceh-30(tm2157); bli-4(e937) unc-37(e262); smIs26; him-5(e1490); ceh-30(tm2157); smIs26; him-5(e1490); ceh-30(tm272); and bli-4(e937) unc-37(e262); smIs26; him-5(e1490); ceh-30(tm272).*

n indicates the number of presumptive CEMs scored.

elevated β -galactosidase activity (Fig. 5B). In contrast, yeast strains expressing GAL4-AD-CEH-30(F42E)/GAL-DB-UNC-37 or GAL4-DB-CEH-30(F42E)/GAL-AD-UNC-37 showed the opposite phenotype in each assay (Fig. 5B). These biochemical data indicate that CEH-30 and UNC-37 can physically interact through the eh1/FIL motif in CEH-30 and that CEH-30 inhibits CEM apoptosis in males by recruiting the transcriptional repressor UNC-37 to the promoter of its target gene(s).

Discussion

In a genetic screen for factors that regulate the sexually dimorphic apoptosis of CEM neurons, we identified *ceh-30*, a BarH homeodomain-encoding gene, as a key inhibitor of CEM cell death. Loss-of-function mutations in *ceh-30* lead to ectopic CEM death in males. Genetic epistasis analysis indicates that *ceh-30* acts downstream from the terminal sex determination factor *tra-1*, upstream of, or in parallel to, the cell death initiator *egl-1*, and probably upstream of the caspase *ced-3* to control the sex-specific death of the CEM neurons (Table 2). Therefore *ceh-30* defines a critical checkpoint that interprets sex specification signals and translates them into an appropriate life versus death response, probably by regulating the expression of a key cell death gene.

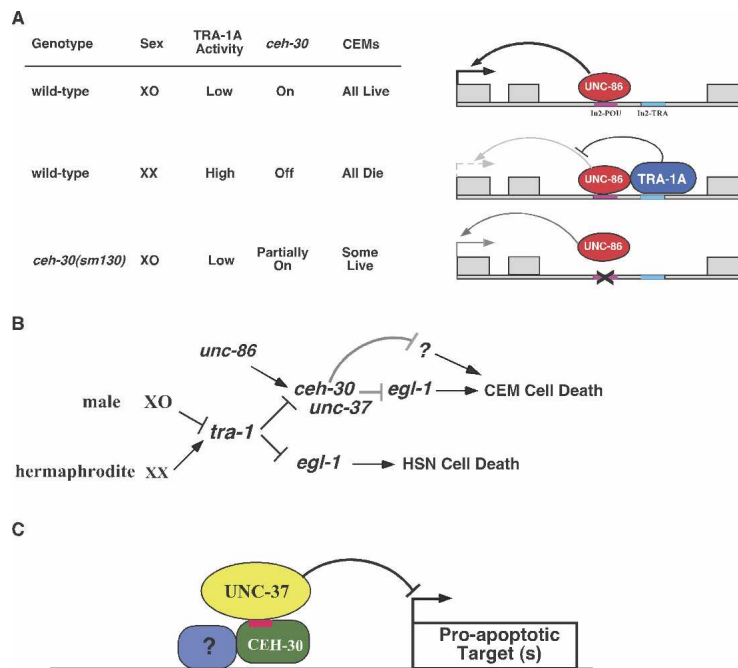
BarH homeodomain proteins were first identified in *Drosophila* due to the “Bar” mutant phenotype and later in several vertebrate species including fish, amphibians, and mammals. They are expressed primarily in the central nervous system, where they are implicated in regulating neuronal cell fates, migration, and survival (Reig et al. 2007). In *C. elegans*, *ceh-30* appears to play a similar role in controlling the cell fate of the chemosensory CEM neurons, promoting the survival of CEMs in males.

Interestingly, deletion of the mouse *Barhl1* gene, which encodes a BarH homeodomain protein, leads to progressive degeneration of cochlear hair cells and loss of many neurons from the zonal layer of the superior colliculus due to increased apoptosis in these regions (Li et al. 2002; Li and Xiang 2006). These observations suggest that BarH homeodomain proteins may play a conserved role in promoting the survival of sensory neurons.

Analysis of two *ceh-30* alleles (*sm130* and *tm272*) reveals a *cis*-regulatory element within the second intron of the *ceh-30* gene that is critical for the survival of CEMs in males. This *cis*-element, In2-POU, resembles a conserved binding site for POU-type homeodomain proteins. Alteration or removal of this In2-POU site by the *sm130* mutation or the *tm272* deletion causes ectopic death of CEMs in males, underscoring its importance for CEM survival. Interestingly, from the same CEM mutant screen, we identified a loss-of-function mutation (*sm117*) in the *unc-86* gene that causes complete absence of CEM neurons. Given that *unc-86* encodes a POU-type homeodomain protein and is expressed in CEM neurons (Finney et al. 1988; Finney and Ruvkun 1990), UNC-86 is an excellent candidate for an In2-POU-binding factor. Indeed, UNC-86 binds In2-POU *in vitro*, and the *sm130* mutation reduces the binding of UNC-86 to the In2-POU site in a manner consistent with the weak reduction of CEM survival observed in *sm130* males. Moreover, the mouse ortholog of UNC-86, Brn3c, also functions genetically upstream of *Barhl1*, a CEH-30 homolog, to specify terminal differentiation and survival of mechanosensory hair cells (Li et al. 2002). These findings suggest that UNC-86/Brn3c and CEH-30/Barhl1 may define an evolutionarily conserved pathway controlling the survival of sensory neurons.

How then does *ceh-30* regulate sex-specific death of CEMs? Interestingly, a conserved TRA-1A-binding site (In2-TRA) is found 55 bp downstream from In2-POU and is altered in several gain-of-function *ceh-30* mutants where CEMs improperly survive in hermaphrodites (H.T. Schwartz and H.R. Horvitz, pers. comm.), suggesting that In2-TRA may control sex-specific death of CEMs in response to the activity of TRA-1A. Indeed, we found that TRA-1A binds loosely to In2-TRA *in vitro* by itself and the binding of TRA-1A to this site is significantly enhanced in the presence of UNC-86, suggesting that UNC-86 interacts with TRA-1 to stabilize TRA-1A binding to intron 2. TRA-1A acts globally in the soma of the hermaphrodite to specify sexual fate. In the developing hermaphrodite, high levels of TRA-1A expression in CEMs presumably would result in binding of TRA-1A to intron 2 of the *ceh-30* gene, which would interfere with or block the transcriptional activation of *ceh-30* by UNC-86 that also binds intron 2. On the other hand, low or no expression of TRA-1A in developing males would allow UNC-86 to activate *ceh-30* expression without interference from TRA-1A and thus promote CEM survival (Fig. 6A). It is worth noting that TRA-1A has also been shown to regulate the sexually dimorphic apoptosis of HSNs by binding to a *cis*-element in the *egl-1* gene and negatively regulating the expression of *egl-1* in hermaph-

Figure 6. Transcriptional regulation of sex-specific CEM cell death. (A) UNC-86, a transcription activator, and TRA-1A, a transcription repressor, interact at intron 2 of the *ceh-30* gene to regulate the expression of *ceh-30*, leading to the proper cell death specification of CEMs. XO indicates males, and XX indicates hermaphrodites. (B) Genetic pathways regulating sexually dimorphic cell deaths in *C. elegans*. *ceh-30* is the key gene that interprets the sex determination signal transduced by *tra-1* and the survival signal specified by *unc-86* to turn off or on programmed cell death in CEMs along with *unc-37*, probably through suppressing the expression of *egl-1* and/or another proapoptotic gene. In HSNs, TRA-1A directly inhibits the expression of *egl-1*. (C) Formation of a CEH-30/UNC-37 repressosome at the promoter of a proapoptotic target gene. A CEM-specific transcription factor that binds to the promoter may recruit the UNC-37/Groucho transcription repressor through CEH-30 via its eh1/FIL motif (in red).



rodite HSNs (Conradt and Horvitz 1999). Therefore, TRA-1A controls appropriate sexually dimorphic apoptosis of HSNs and CEMs by repressing the expression of two downstream target genes with contrasting apoptotic functions: the death-initiating *egl-1* gene in HSNs and the death-inhibiting *ceh-30* gene in CEMs (Fig. 6B).

Given the importance of homeodomain proteins in cell fate determination, it is very surprising that the *ceh-30* deletion allele (*tm2157*), which removes the CEH-30 homeodomain, has only a weak effect on the survival of male CEMs. This finding suggests that the homeodomain of CEH-30 is largely dispensable for its CEM death protective function and that the N terminus of CEH-30 is sufficient to confer most of its death protective activity. The N terminus of CEH-30 contains a conserved eh1/FIL motif that has been shown to mediate protein-protein interaction and recruitment of the general transcriptional repressor *Groucho* (Smith and Jaynes 1996; Choi et al. 1999; Jimenez et al. 1999; Bae et al. 2003). Indeed, this eh1/FIL motif in CEH-30 mediates interaction of CEH-30 with the *C. elegans* *Groucho* ortholog UNC-37 (Fig. 5B), and a mutation (F42E) that alters the conserved and critical Phe residue in this eh1/FIL motif disrupts the binding of CEH-30 to UNC-37 and the death protective function of CEH-30 in male CEMs. Furthermore, a mutation (*e262*) that reduces the activity of UNC-37 not only can cause ectopic male CEM death on its own but also can synergize with the *ceh-30(tm2157)* mutation to cause ectopic death of most male CEMs, suggesting that CEH-30 and UNC-37 act together to inhibit CEM cell death in males. It is interesting that both *ceh-30* and *unc-37* are widely expressed in *C. elegans* embryos (Fig. 4A; Winnier et al. 1999), yet *ceh-30* specifically represses cell death only in male CEMs. This observation and the finding that the CEH-30 homeodo-

main is largely dispensable for its death inhibitory activity in CEMs suggest that CEH-30 may act to bridge the repressosome formation of UNC-37/*Groucho* with a CEM-specific transcription factor that directs repressosome binding to the promoter of a critical proapoptotic target gene and thus represses CEM cell death in males (Fig. 6C). In total, our study reveals a delicate transcriptional regulatory cascade that responds to appropriate sexual signals to repress or activate apoptosis in a sex-specific and cell-type-specific manner, thereby generating appropriate male-specific neurons (Fig. 6B).

Materials and methods

Strains

C. elegans strains were maintained at 20°C, unless otherwise noted. The N2 (Bristol) strain was the standard wild-type strain. For single-nucleotide polymorphism (SNP) mapping, the Hawaiian strain CB4856 was used. The alleles that were used in this study are LGI: *ceh-6(mg60)*, *unc-37(e262)*, *bli-4(e937)*, *dpy-5(e61)*, *unc-13(e450)*, and *hT2(I;III)*; LGII: *smIs23*; LGIII: *tra-1(e1099)*, *unc-86(sm117, n845, e1416, e1507)*, and *dpy-18(e364)*; LGIV: *smIs26*, *him-8(e1489)*, and *ced-3(n717, n2433)*; LGV: *egl-1(n1084n3082)*, *him-5(e1490)*, and *unc-76(e911)*; and LGX: *ceh-30(sm130, tm272, tm2157)*, *lon-2(e678)*, *unc-2(e55)*, *dpy-3(e27)*, *ceh-18(mg57)*, and *smIs54*.

EMS mutagenesis

EMS mutagenesis was carried out as described previously (Brenner 1974). Briefly, mixed-stage worms were exposed to 47 mM EMS for 4 h with agitation. The F₁ progeny of mutagenized animals were cloned out, and their F₂ progeny were screened for loss of CEMs in males and improper survival in hermaphrodites.

Mapping of *sm130*

sm130 was mapped to LGX by crossing *smIs23*; *him-5(e1490)*; *sm130* males into *smIs23*; *him-5(e1490)*; *lon-2(e678)* hermaph-

rodites, picking non-Lon cross progeny, and scoring the male progeny of Lon F₂ animals for the CEM defect. Further three-factor mapping was done by crossing *smIs23*; *him-5(e1490)*; *sm130* males into *unc-2(e55) lon-2(e678)* hermaphrodites and scoring Lon non-Unc or Unc non-Lon F₂ progeny. The location of *sm130* was refined by consecutive SNP mapping using several SNP markers: *snp_F53B3*, *snp_F52E4*, *snp_K09C4*, *snp_F11D5*, and *snp_C02F12* (Wicks et al. 2001). To obtain X-chromosome recombinants between *sm130* animals and the SNP-rich Hawaiian strain (CB4856), *smIs26* males carrying the X chromosome derived from CB4856 were crossed into *smIs26*; *him-5(e1490)*; *dpy-3(e27) sm130 lon-2(e678)* hermaphrodites. Lon non-Dpy homozygous recombinants were analyzed for the presence of *sm130* by scoring for CEM loss in males, and SNPs were genotyped by PCR amplification and subsequent restriction enzyme digestions. This SNP mapping strategy placed *sm130* between two SNPs on Cosmids F53B3 and F52E4.

Scoring of CEMs and HSNs

The presence of CEMs was scored in L4 larvae and adults using the integrated *P_{pkd-2}gfp* reporter, *smIs23*, or *smIs26*. CEMs can also be scored in L4 larvae based on their cell morphology using Nomarski optics. The percentage of surviving CEMs was calculated by dividing the total number of CEMs observed by the maximum possible number of CEMs [(the number of CEMs observed)/(4 × the number of animals scored)]. The presence of HSNs was scored in L4 larvae using *smIs26*, which also carries *P_{trph-1}gfp* that directs GFP expression in HSNs (Sze et al. 2000) or by Nomarski optics in L1 or L4 larvae based on HSN cell morphology.

Molecular biology

C33D12 was digested with *NheI*, and the resulting 9496-bp *ceh-30*-containing fragment was cloned into the pBluescript SK(-) vector (Stratagene) via its *XbaI* site. This construct was then digested with *SacI* to remove a 1786-bp fragment, leaving a 7710-bp C33D12 *NheI*-*SacI* fragment. *P_{ceh-30}ceh-30::gfp* was constructed by PCR-amplifying the *ceh-30* promoter and ORF from the first C33D12 9.5-kb construct using a *ceh-30*-specific primer 5'-GCTCTAGATTCTGAGTTGCTGGAAACATCC-3' (contains an *XbaI* site) and a primer from the vector (T7: 5'-AATACGACTCACTATAG-3'). The resulting 5.8-kb PCR product was subcloned into pPD95.77 via its *PstI* and *XbaI* sites. Full-length *ceh-30* cDNA was generated by RT-PCR using primers 5'-GCGATATCCTATTCTGAGTTGCTGGAAACATCC-3' (contains an *EcoRV* site) and 5'-GCGCTAGCATGTCACTTCTCGACCCTCGGC-3' (contains an *NheI* site). The amplified cDNA fragment was subcloned into pSL1190 (Amersham Biosciences) via its *NheI* and *EcoRV* sites. *P_{ceh-30}ceh-30(Δintron2)::gfp* was generated by PCR amplification of the *ceh-30* cDNA using primers 5'-GCGCTAGCATGTCACTTCTCGACCCTCGGC-3' and 5'-GCTCTAGATTCTGAGTTGCTGGAAACATCC-3' (contains an *XbaI* site) followed by digestion with *KpnI* (within exon 2) and *XbaI*. The resulting *KpnI*-*XbaI* fragment was subcloned into *P_{ceh-30}ceh-30::gfp* that had been partially digested with *KpnI* and *XbaI*. *P_{ceh-30}ceh-30(ΔIn2-POU)::gfp*, *P_{ceh-30}ceh-30(1-92)::gfp*, and *P_{ceh-30}ceh-30(F42E)::gfp* constructs were generated using Stratagene's QuikChange Site-Directed Mutagenesis Kit with the following primers: 5'-AATTGTGCCTCTTTTGAATGTATCCACAATCTGGAAACCTTTTCAAACGT-3' and 5'-GACGTTTTGAAAGGTTTTCCAGATTGTGGATACATTCAAAGAGGCACAAAT-3' for making *P_{ceh-30}ceh-30(ΔIn2-POU)::gfp*, 5'-CCAGGCAGTTGCTAAATATCTAGAAAGCAAGAACC-3' and 5'-GGTCTTGCC

TTTCTAGATTATTAGCAACTGCCTGG-3' for making *P_{ceh-30}ceh-30(1-92)::gfp*, and 5'-CAAAATTCGTCCTTCTTCTTCA GAGCGGATATCTGACATTCTCGAGCAATCC-3' and 5'-GGGATTGCTCGAGAATGTCAGATATCCGCTCTGAAGAA GAAGACGAAATTTT-3' for *P_{ceh-30}ceh-30(F42E)::gfp*.

Transgenic animals

Germline transformation was performed as described previously (Mello et al. 1991). Cosmid DNA, rescuing constructs, and GFP fusion reporters (20 ng/μL each) were injected into animals with the appropriate genetic background using *P_{sur-5}RFP* as a coinjection marker, which directs expression of red fluorescent protein (RFP) in every cell of *C. elegans* larvae (Gu et al. 1998).

Expression and purification of proteins

Full-length UNC-86 was purified as described previously (Xue et al. 1993). Full-length *tra-1* cDNA was subcloned into the pET-30b vector (Novagen) via its *KpnI* and *NotI* sites. BL21(DE3) Lys S bacterial cells were transformed with the appropriate expression vector, cultured to an OD₆₀₀ of 0.4–0.5, induced to express the protein with 1 mM IPTG for 1 h at room temperature. Cells were then lysed by sonication in buffer A (10% glycerol, 20 mM Tris-HCl at pH 7.5, 200 mM NaCl, 5 mM imidazole, 0.1% NP-40, and 1 mM PMSF). The soluble fraction was incubated with Ni²⁺-NTA-agarose beads for 4 h at 4°C. The beads were washed with 50 vol of wash buffer (buffer A with 60 mM imidazole and no PMSF). Bound proteins were eluted with buffer A containing 250 mM imidazole but no PMSF.

Gel mobility shift assays

For gel mobility shift analysis, oligonucleotides or PCR fragments were labeled with ³²P-ATP using T4 polynucleotide kinase. Purified proteins (25 ng of GST::UNC-86 and/or 5 ng of TRA-1::His₆) were preincubated with 0.05 mg/mL poly(dG-dC) in a buffer containing 4% glycerol, 1 mM MgCl₂, 0.5 mM EDTA, 0.5 mM DTT, 50 mM NaCl, and 10 mM Tris-HCl (pH 7.5) for 10 min at room temperature. Labeled DNA was then added and incubated for another 20 min at room temperature. Samples were resolved on 4% or 8% native polyacrylamide gels at 4°C. Gels were dried and exposed to PhosphorImaging screens overnight. Quantification of gel shift bands was done with the ImageQuant software package (GE Healthcare). Antibody supershift experiments were performed as described previously (Xue et al. 1993).

Yeast two hybrid

Yeast two-hybrid assays were performed using the ProQuest Two-Hybrid System with Gateway Technology (Invitrogen). GAL4 fusion proteins were generated using the following PCR primers for *unc-37* and *ceh-30*, respectively: 5'-GGGGACAA GTTTGTACAAAAAGCAGGCTTCATGAAGGCATCGTA TCTGG-3', 5'-GGGGACCACTTTGTACAAGAAAGCTGGG TCTTAATATTCAACTGCATAGA-3', 5'-GGGGACAAGTTT GTCAAAAAAGCAGGCTTCATGTCACTTCTCGACCCTC GG-3', and 5'-GGGGACCACTTTGTACAAGAAAGCTGGG TCCTATTCTGAGTTGCTGGAAACATCC-3'. Yeast growth was assayed on synthetic complete SC-Leu-Trp plates lacking histidine but containing 10 mM 3AT, or lacking uracil while containing 0.2% 5FOA. The X-gal staining and the liquid β-galactosidase assay using chlorophenol-red-β-D-galactopyrano-

side (CPRG) as a substrate were performed as described in the ProQuest manual.

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